

11-15-00

PTO/SB/05 (4-98)

Approved for use through 09/30/2000 OMB 0651-0032
Patent and Trademark Office U.S. DEPARTMENT OF COMMERCE

Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it displays a valid OMB control number

**UTILITY
PATENT APPLICATION
TRANSMITTAL**

(Only for new nonprovisional applications under 37 C.F.R. § 1.53(b))

Attorney Docket No. SD-8310

First Inventor or Application Identifier MESCHER

Title LITHOGRAPHY PROCESS FOR PATTERNING HgI₂ PHOTONIC
DEVICES

Express Mail Label No. EK770676429US

APPLICATION ELEMENTS

See MPEP chapter 600 concerning utility patent application contents.

1. ☒ *Fee Transmittal Form (e.g., PTO/SB/17)
(Submit an original and duplicate for fee processing)
2. ☒ Specification [Total Pages **21**]
(preferred arrangement set forth below)
- Descriptive title of the invention
- Cross References to Related Applications
- Statement Regarding Fed sponsored R & D
- Reference to Microfiche Appendix
- Background of the invention
- Brief Summary of the invention
- Brief Description of the Drawings (if filed)
- Detailed Description
- Claim(s)
- Abstract of the Disclosure
3. ☒ Drawing(s) (35 U.S.C. 113) [Total Sheets **2**]
4. ☒ Oath or Declaration [Total Pages **4**]
a. ☒ Newly executed (original or copy)
b. ☐ Copy from a prior application (37 C.F.R. § 1.63(d))
(for continuation/divisional with Box 16 completed)
i. ☐ **DELETION OF INVENTOR(S)**
Signed statement attached deleting
inventor(s) named in the prior application,
see 37 C.F.R. §§ 1.63(d)(2) and 1.33(b).

*NOTE FOR ITEMS 1 & 13 IN ORDER TO BE ENTITLED TO PAY SMALL ENTITY
FEES, A SMALL ENTITY STATEMENT IS REQUIRED (37 C.F.R. § 1.27), EXCEPT IF
ONE FILED IN A PRIOR APPLICATION IS RELIED UPON (37 C.F.R. § 1.29)

ADDRESS TO: Box Patent Application
Assistant Commissioner for Patents
Washington, DC 20231

5. ☐ Microfiche Computer Program (Appendix)
6. ☐ Nucleotide and/or Amino Acid Sequence Submission
(if applicable, all necessary)
a. ☐ Computer Readable Copy
b. ☐ Paper Copy (identical to computer copy)
c. ☐ Statement verifying identity of above copies

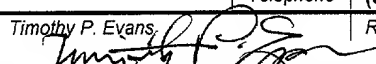
ACCOMPANYING APPLICATION PARTS

7. ☐ Assignment Papers (cover sheet & document(s))
8. ☐ 37 C.F.R. § 3.73(b) Statement
(when there is an assignee)
9. ☐ English Translation Document ☐ Power of
(if applicable) Attorney
10. ☐ Information Disclosure ☐ Copies of IDS
Statement (IDS)/PTO-1449 Citations
11. ☐ Preliminary Amendment
12. ☒ Return Receipt Postcard (MPEP 503)
(Should be specifically itemized)
13. ☐ *Small Entity ☐ Statement filed in prior application,
Statement(s) Status still proper and desired
(PTO/SB/09/12)
14. ☐ Certified Copy of Priority Document(s)
(if foreign priority is claimed)
15. ☐ Other: _____

16. If a CONTINUING APPLICATION, check appropriate box, and supply the requisite information below and in a preliminary amendment☐ Continuation ☐ Divisional ☐ Continuation-in-part (CIP) of prior application No. _____ / _____
Prior application information. Examiner _____ Group / Art Unit. _____

For CONTINUATION or DIVISIONAL APPS only: The entire disclosure of the prior application, from which an oath or declaration is supplied under Box 4b, is considered a part of the disclosure of the accompanying continuation or divisional application and is hereby incorporated by reference. The incorporation can only be relied upon when a portion has been inadvertently omitted from the submitted application parts.

17. CORESPONDENCE ADDRESS☒ Customer Number or Bar Code Label **0215868** or ☐ Correspondence address below
(Insert Customer No. or Attach bar code label here)

Name	Timothy Evans				
	MS 9031				
Address	Sandia National Laboratories				
	7011 East Avenue				
City	Livermore	State	CA	Zip Code	94550
Country	USA	Telephone	(925) 294-3690	Fax	(925) 294-3389
Name (Print/Type)	Timothy P. Evans		Registration No. (Attorney/Agent)		
Signature			41,013		Date 11/13/00

Burden Hour Statement: This form is estimated to take 0.2 hours to complete. Time will vary depending upon the needs of the individual case. Any comments on the amount of time you are required to complete this form should be sent to the Chief Information Officer, Patent and Trademark Office, Washington, DC 20231. DO NOT SEND FILES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Assistant Commissioner for Patents, Box Patent Application, Washington, DC 20231

FEE TRANSMITTAL FOR FY 2000

Patent fees are subject to annual revision.
Small Entity payments must be supported by a small entity statement.
Otherwise large entity fees must be paid See Forms PTO/SB/09-12

TOTAL AMOUNT OF PAYMENT (\$)**980.00**

Complete if Known

Application Number	not assigned
Filing Date	11/13/2000
First Named Inventor	MESCHER
Examiner Name	not assigned
Group / Art Unit	not assigned
Attorney Docket No.	SD-8310

METHOD OF PAYMENT (check one)

The Commissioner is hereby authorized to charge
1. ☒ indicated fees and credit any over payments to:

Deposit Account Number **50-0583**

Deposit Account Name **SNL by KCO**

☐ Charge Any Additional
Fee Required Under
37 CFR 1.16 and 1.17

2. Payment Enclosed:

☐ Check ☐ Money Order ☐ Other

FEE CALCULATION

1. BASIC FILING FEE

Large Fee Code	Entity Fee (\$)	Small Fee Code	Entity Fee (\$)	Fee Description	Fee Paid
101	690	201	380	Utility filing fee	710.
106	310	206	155	Design filing fee	
107	480	207	240	Plant filing fee	
108	690	208	380	Reissue filing fee	
114	150	214	75	Provisional filing fee	

SUBTOTAL (1) **\$ 710.00**

2. EXTRA CLAIM FEES

	Extra Claims	Fee from below	Fee Paid
Total Claims 35	-20**= 15	X 18.	= 270.00
Independent Claims 1	- 3**= 0	X 80.	= 0.00
Multiple Dependent			=

**or number previously paid, if greater, For Reissues, see below

Large Fee Code	Entity Fee (\$)	Small Fee Code	Entity Fee (\$)	Fee Description
103	18	203	9	Claims in excess of 20
102	80	202	40	Independent claims in excess of 3
104	270	204	135	Multiple dependent claim, if not paid
109	80	209	40	**Reissue independent claims over original patent
110	18	210	9	**Reissue claims in excess of 20 and over original patent

SUBTOTAL (2) **(\$270.00)**

FEE CALCULATION (continued)

3. ADDITIONAL FEES

Large Fee Code	Entity Fee (\$)	Small Fee Code	Entity Fee (\$)	Fee Description	Fee Paid
105	130	205	65	Surcharge - late filing fee or oath	
127	50	227	25	Surcharge - late provisional filing fee or cover sheet	
139	130	139	130	Non-English specification	
147	2,520	147	2,520	For filing a request for reexamination	
112	920*	112	920*	Requesting publication of SIR prior to Examiner action	
113	1,840	11	1,840	Requesting publication of SIR after Examiner	
115	110	215	55	Extension for reply within first month	
116	390	216	195	Extension for reply within second month	
117	890	217	445	Extension for reply within third month	
118	1,390	218	695	Extension for reply within fourth month	
128	1,890	228	945	Extension for reply within fifth month	
119	310	219	155	Notice of Appeal	
120	310	220	155	Filing a brief in support of an appeal	
121	270	221	135	Request for oral hearing	
138	1,510	138	1,510	Petition to institute a public use proceeding	
140	110	240	55	Petition to revive - unavoidable	
141	1,240	241	620	Petition to revive - unintentional	
142	1,240	242	620	Utility issue fee (or reissue)	
143	440	243	220	Design issue fee	
144	600	244	300	Plant issue fee	
122	130	122	130	Petitions to the Commissioner	
123	50	123	50	Petitions related to provisional applications	
126	240	126	240	Submission of Information Disclosure Stmt	
581	40	581	40	Recording each patent assignment per property (times number of properties)	
146	710	246	355	Filing a submission after final refection (37 CFR 1.129(a))	
149	710	249	355	For each additional invention to be examined (37 CFR 1.129(b))	
Other fee (specify) _____					
Other fee (specify) _____					

*Reduced by Basic Filing Fee Paid

SUBTOTAL (3) **(\$)**

SUBMITTED BY

Name (Printer/Type) **Timothy P. Evans**

Signature 

Complete (if applicable)

Reg Number (Attorney/Agent) **41,013**

Telephone **(925) 294-3690**

Date **11/13/00**

WARNING:

Information on this form may become public. Credit card information should not be
included on this form. Provide credit card information and authorization on PTO-2038

CERTIFICATION UNDER 37 CFR 1.10

I hereby certify that this New Application Transmittal and the documents referred to as enclosed therein are being deposited with the U. S. Postal Service on **NOVEMBER 13, 2000**, in an envelope as "Express Mail Post Office to Addressee" Mailing Label Number EK770676429US, addressed to: Box Patent Application, Assistant Commissioner for Patents, Washington, D.C. 20231.

Date of Deposit: 11/13/00

T P Evans
Person Making Deposit
[Signature]
Signature

"Express Mail" Mailing Label No. EK770676429US

**LITHOGRAPHY PROCESS FOR PATTERNING HgI₂
PHOTONIC DEVICES**

INVENTORS:

Mark J. MESCHER
Auburndale, MA 02466

Ralph B. JAMES
Livermore, CA 94550

Haim HERMON
Jerusalem,
State of Israel

CITIZENS OF THE UNITED STATES OF AMERICA

LITHOGRAPHY PROCESS FOR PATTERNING HgI₂ PHOTONIC DEVICES

The United States Government has rights in this invention pursuant to Contract No. DE-AC04-94AL85000 between the United States Department of Energy and Sandia Corporation for the operation of Sandia National Laboratories.

BACKGROUND OF THE INVENTION

Field of the Invention

The present invention relates to methods for obtaining improved spatial and energy resolution in room temperature HgI₂ radiation detectors, and more specifically, it relates to a method for lithographically patterning HgI₂ crystals.

Description of Related Art

Semiconductors materials exhibit a band gap between their valence and conduction bands of typically a few eV. Because this energy gap is so low, as the temperature of the crystal is increased, electrons are thermally excited and easily move from the valance band to the conduction band. The electrical properties of these materials, therefore, are effected not only by the movement of electrons into the conduction band but also by the formation of vacant sites or "holes" in the valence bands left behind by the departing electrons. Both can conduct current.

Holes also may be created by the interaction of energetic radiation, such as X-rays, gamma rays, and the like, with intrinsic semiconductors and, therefore, one should be able to use these materials as detectors for measuring high-energy radiation. In fact, high-resistivity semiconductor radiation detectors are widely used for detecting ionizing radiation due to their ability to operate at room temperature, their small size and durability. Such detectors are used in a wide variety of applications, including medical diagnostic imaging, nuclear waste monitoring, industrial process monitoring, and space astronomy. Ionizing radiation includes both particulate radiation, such as alpha or beta particles, and electromagnetic radiation, such as gamma or x rays.

If all the electrons and holes generated by the ionizing radiation reach their respective electrodes (i.e., the electrons reach the anode and the holes reach the cathode), the output charge signal will exactly equal the charge from the energy deposited within the crystal by the radiation. Because the deposited charge is directly proportional to the energy of the ionizing radiation, the semiconductor detector provides a means for measuring the energy of the ionizing radiation.

Room temperature detectors, however, suffer from a serious drawback. Because of limitation in the transport properties of the bulk semiconductor crystal, some of the electrons and, more particularly, some holes are generally lost by being trapped as they move toward the respective electrodes under the influence of the external electrical field. This is particularly evident for semiconductors wherein the transport properties of one carrier type (e.g., electrons) are much better than those of another type (in this example the "holes"). Under such circumstance, therefore, the amplitude of the output charge signal becomes dependent on the position within the crystal at which the ionizing radiation is absorbed. Generally speaking, the amplitude is less than the charge deposited by the ionizing radiation and results in a corresponding reduction of energy measurement accuracy, poor resolution, and reduced peak efficiency. This loss (or trapping) of charge in a radiation detector results in distorted and asymmetrical spectral peak shapes known as "hole tailing" or "hole trapping."

The inability to eliminate "hole" drift current is a major impediment for the use of room temperature semiconductors as detectors. Gamma-ray spectroscopy is particularly encumbered because pulse height spectra produced by these devices are distorted by this process. Mono-energetic gamma rays produce charge signal responses of different pulse height because the total combined distance drifted by the electrons and holes is dependent on the position of gamma-ray interaction. This phenomenon is well known in the prior art and has been described by many researchers. It is widely understood to be the major

deficiency limiting the effectiveness of room temperature semi-conductor materials.

Due to the deleterious effects of hole-trapping in semiconductor detectors, much effort has gone into attempting to solve this problem. U.S. Pat. Nos. 4,253,023 and 4,996,432 recognized the problem and proposed early remedies. The first of these included a method to de-convolute the contribution of the electron motion from the acquired signal. The second approach proposes a method relying upon use of a thick crystal and a crystal orientation placing the detector anode surface facing the source of radiation thereby reducing the positional dependence of the radiation interaction with the crystal and restricting it only to that part of the crystal immediately behind the anode. Neither of these approaches directly addresses the problem of eliminating hole-trapping.

U.S Pat. No. 5,677,539 provides another approach and a comprehensive review of much of the pertinent prior art. A particularly relevant approach, described therein, employs an anode patterned into an interleaving grid structure, with the cathode remaining planar. (See, e. g., P. N. Luke, "Unipolar Charge Sensing with Coplanar Electrodes- Application to Semiconductor Detectors," IEEE Tran. Nucl. Science, vol. 42, No. 4, at pages 207-213 (1995)). In this approach, one set of anode grids is maintained at a slightly higher voltage than the other. A train of signal conditioning electronics is connected to each set of grids, and the difference between the outputs from these trains constitutes the final output signal. With this arrangement, when the charge cloud is far from the grids, the difference-signal between the grid outputs is zero. As the cloud approaches the grids, the induced charge on one grid rises rapidly, while the charge induced in the other grid drops rapidly. The difference signal is then a measure of the full charge in the electron cloud, independent of the position of the ionizing event.

This approach, however, also suffers from various drawbacks. First, the grid structure is relatively complex and would be difficult, if not impossible, to use in detector arrays. Second, the grids require two separate amplifying chains, plus

a difference amplifier, adding significantly to the complexity and cost of manufacture. This circuitry also would be difficult to implement in the multichannel type integrated circuits needed in detector array structures.

A relatively simpler structure is a variation on a technique devised by Frisch for use in gas detectors. The hole-trapping phenomenon observed in semiconductor detectors is analogous to the trapping behavior of positive ions in gas detectors. Frisch proposed, and later developed, detectors that contained a grid of conductive wires between the two electrodes of a conventional gas detector.

In the Frisch grid type detector, the signal is measured between the anode and the grid. The negative charge carriers, electrons in the case of a semiconductor detector, usually drift all of the way to the anode. Thus, any radiation interactions occurring between the cathode and the grid will produce electrons that drift past the grid and on to the anode. When the electrons drift across the gap between the grid and the anode, they will induce a current that is due solely to the motion of electrons. The current induced by positively charge carriers ("holes" in the case of semiconductors) traveling in the opposite direction is shielded by the grid. The main advantage, then, of the Frisch grid is that the signal produced from the device is independent of the position of interaction between the cathode and grid (completely solving the hole trapping problem in the cathode to grid region).

An approach to reducing hole-trapping in a semiconductor detector using a modified Frisch grid approach has been proposed by D. S. McGregor.

McGregor's scheme, described as an "etch trench" device, entailed building a Frisch Grid device on a semiconductor crystal and placing electrodes at the bottom of these trenches. The primary disadvantage of McGregor's etch-trench design is that it is difficult to execute. In other words, it might be very difficult to produce the requisite trenches, particularly since the fabrication technology for room temperature semiconductor materials is not well developed at this time.

Therefore, while a Frisch-grid type device design would work very well in a room

temperature semiconductor detector, it suffers from the fundamental drawback of being difficult to construct. Such a device, owing to the difficulty of placing a conductive grid inside a semiconductor crystal would be inherently complex and expensive.

5 A third approach is discussed in U.S. Pat. No. 5,677,539. This invention takes advantage of the principle that a significant reduction in tailing in a semiconductor detector can be attained by a novel arrangement of electrodes that share induced charge from ionizing events in the detector, that properly shape the electric field, and that focus charge collection toward a small electrode.

10 The detector includes three electrodes formed on the surface of a semiconductor crystal. The crystal has a plurality of sides; it preferably has a thickness of at least about 0.5 mm and is preferably formed from a semiconductor material having a higher mobility-lifetime product for one electronic charge carrier compared to the other. The first electrode is a bias
15 electrode, which preferably covers the entire surface of one side of the crystal. At least one signal electrode having a small area is preferably formed on the opposing side of the crystal from the bias electrode. A control electrode is preferably disposed on the same side containing the signal electrode.

20 In particular, the control electrode is formed on the same side of the semiconductor crystal as the signal electrode (anode), and the bias electrode (cathode) covers substantially the entire surface of the opposite side of the crystal. The semiconductor crystal is formed from CdZnTe or CdTe. In the simplest configuration, the anode is a small contact point located near the center of the electron-charge-collection side of the crystal. The anode is coupled to
25 ground through a large-value resistor and to external signal circuitry. The cathode is coupled to a voltage source that maintains the cathode at a negative voltage level relative to the anode. Preferably, the control electrode is much larger in area than the anode and forms a single ring surrounding the anode. The control electrode is maintained at a voltage level that is negative with respect to
30 the anode, but generally not more negative than the cathode.

This approach relies on a geometric construction of the various electrodes to shape the electric field within the crystal in order to direct and accelerate the more mobile of the charge carriers toward the appropriate collecting electrode. By operating in this manner the less mobile of the charge carriers, those that are

5 "trapped", arrive typically too late to contribute to the analysis. However, while much reduced, some small percentage of these carriers still do contribute to the signal.

The performance of gamma-ray and x-ray spectrometers fabricated from Hgl₂ is substantially limited by hole trapping. The ability to pattern arbitrarily

10 sized and positioned electrodes allows for enhanced performance via the so-called "electron-only" behavior, which substantially increases detector energy resolution. Hole trapping distorts the current pulse generated by the absorption of an x- or gamma-ray, which reduces the ability of the detectors to spectrally resolve the unique radiological emissions of a wide variety of isotopes.

15 Degradation in the capability of the spectrometers to resolve energies of x-rays and gamma-rays reduces the marketability of the devices.

For imaging devices, the spatial resolution is limited by the pixel size of each individual detector element. It is desirable that pixel size be significantly decreased as compared to the current method of shadow mask deposition. The

20 ability to pattern substantially denser arrays of smaller pixels will improve the spatial resolution of these imaging devices.

SUMMARY OF THE INVENTION

It is an object of the present invention to provide a method using a photolithographic process for forming patterns on Hgl₂ surfaces.

25 It is another object of the present invention to provide a method for smoothing Hgl₂ surfaces.

Still another object of the present invention is to provide a method for producing trenches in Hgl₂.

It is another object of the present invention to provide a sublimation process for trench etching to produce etched-trench devices with enhanced electron-transport-only behavior.

Another object of the invention is to provide a lithographic process for
5 defining a metal sublimation mask and electrodes to substantially improve device performance by increasing the realizable design space.

These and other objects of the invention will be apparent to those skilled in the art based on the teachings herein.

The invention patterns HgI_2 surfaces with a new photolithographic
10 process. The process is used to define metal sublimation masks and electrodes on HgI_2 . Design space is increased to substantially improve device performance. Techniques for smoothing HgI_2 surfaces and for producing trenches in HgI_2 are provided. A sublimation process is described which produces etched-trench devices with enhanced electron-transport-only behavior.

15 To deposit a continuous Pd (or other metal) film on a given substrate (e.g. mercuric iodide), the film thickness must be about as thick as (or greater than) the surface roughness. Rapid agitation of the etch solution produced smooth surfaces. The invention provides agitation parameters that produce optimal smoothness.

20 Materials used for both electrode and sublimation mask metal layers include palladium, gold, carbon, indium-tin-oxide, chromium, tantalum and platinum. The films are deposited using techniques such as sputtering and electron beam deposition. Although sputtering will generally not cause substrate heating above 50-100°C for plasma powers less than 200 W, it was necessary to
25 cool the substrate to optimize adhesion.

Shadow masks are typically used to pattern HgI_2 because acetone and/or other solvents used to remove the photoresist after patterning attack HgI_2 rapidly. HgI_2 cannot withstand the high temperatures typically required for photoresist baking. Shadow masks, however, are quite limiting in terms of minimum pattern
30 size and freedom of design. Fine line widths of guard electrodes are required for

electron-only operation of HgI_2 spectrometers, and shadow masks are often not suitable for generating them.

The present invention provides a new lithography process that is compatible with HgI_2 . A low-temperature photoresist is employed in the new process. Solvents are not used for dissolving the photoresist after the metal etching step. A high-temperature bake was not used. The palladium etchant was chosen such that a soft-bake was sufficient to make the photoresist a suitable mask for the Pd etch step. Without the hardbake step, the photoresist removal can subsequently be promoted by a flood exposure step after the Pd etching.

Trenches were etched by a sublimation process after the first Pd metallization and patterning steps. In the present invention, sublimation is governed by temperature only and not by gas flow as long as the pressure is kept somewhat below 11 mTorr. In addition to being temperature dependent, the sublimation rate was also found to be dependent on the particular etchant used for the Pd patterning, with an increase in Br content generating higher etch rates and a redder surface. After the sublimation step was completed, the entire surface was metallized again and patterned with collection and grid electrodes.

BRIEF DESCRIPTION OF THE DRAWINGS

FIGURES 1A through 1E show mercuric iodide after various processing steps.

FIGURE 1A shows lithography on Pd to define grid etch pit.

FIGURE 1B shows etching of Pd.

FIGURE 1C shows HgI_2 sublimation etch (300 μm).

FIGURE 1D shows lithography for grid and collecting electrodes.

FIGURE 1E shows mercuric iodide after Pd etching.

FIGURE 2 shows a schematic cross-sectional diagram for an example single-carrier device where the sensor signal comes from the contacts measuring just the electron current.

DETAILED DESCRIPTION OF THE INVENTION

The invention provides a photolithographic process for forming patterns on Hgl₂ surfaces and for defining metal sublimation masks and electrodes to substantially improve device performance by increasing the realizable design space. Techniques for smoothing Hgl₂ surfaces and for producing trenches in Hgl₂ are provided. A sublimation process is described which produces etched-trench devices with enhanced electron-transport-only behavior.

Surface Preparation

An improvement in surface preparation was required in order to be able to deposit a Pd electrode layer (or other material such as gold or platinum) onto mercuric iodide that could be patterned to μm level dimensions. In order to deposit a continuous film on a given substrate, it is necessary that the film thickness be approximately as thick as (or greater than) the surface roughness. Since film thickness is limited to several thousand angstroms due to stress-related delamination problems, the surface roughness should be less than 1000 Å. Experimentation using a standard procedure to prepare aged Hgl₂ samples for processing revealed surface roughnesses considerably larger than this. The standard for etching aged Hgl₂ crystals (which are typically iodine deficient) is typically a 2-5 minute etch in a 5-10% solution of KI in H₂O. This will generate a nearly stoichiometric surface, but with large surface roughness (Table 1). It was found that rapid agitation of the etch solution produced considerably smoother surfaces. It is supposed here that the etching mechanism is transport-limited without agitation, and thus large gradients in the etchant and etch-byproduct concentrations could exist along the surface of the crystal due to non-uniform stoichiometry and thus local etch rate. This could generate surface roughness. If agitation is used to increase the transport rate, these concentration gradients are limited to much smaller distances which should effectively make the etch rate more uniform and thus decrease surface roughness. Table 1 shows surface roughnesses measured by a Dektak profilometer for various etching conditions.

Roughly one order of magnitude decrease in surface roughness can be achieved with the proper agitation level (conditions shown in Table 2).

Table 1. Measured surface roughness for various etching conditions of aged HgI_2 crystals

Sample condition	Surface roughness [\AA]
aged, no etch	4900
acetone, no agitation	2800
1:10 $\text{KI}:\text{H}_2\text{O}$, no agitation	5000
1:10 $\text{KI}:\text{H}_2\text{O}$, slow agitation (100 rpm)	620
1:10 $\text{KI}:\text{H}_2\text{O}$, fast agitation (200 rpm)	320

5 **Table 2.** Conditions used for optimal surface etch treatment.

Etch solution	time	Sample rotation sequence	agitation	rinse	dry
1:10 $\text{KI} / \text{H}_2\text{O}$	30s 4x	$180^\circ, 90^\circ, 180^\circ$	200 rpm (stir bar)	H_2O	N_2

Metal Deposition

Palladium, which is typically used as the electrode material for mercuric iodide, was used for both electrode and sublimation mask metal layers. Other usable materials include gold, carbon, indium-tin-oxide, chromium, tantalum, and platinum. The films were sputtered using a DC magnetron source because of the low temperatures that can be maintained relative to evaporation. Other film deposition techniques, such as electron beam deposition, may be used. Two aspects of the sputtering process are of primary importance. First, a low base pressure of the system before deposition is important for good film adhesion (adhesion was qualitatively determined by fractional metal coverage after the lithography step). High vacuum at room temperature induces enough sublimation that good surface stoichiometry is insured. Second, substrate

temperature must be kept low to obtain good film adhesion. Although sputtering will generally not cause substrate heating above 50-100°C for plasma powers less than 200 W, it was necessary to cool the substrate to optimize adhesion. A two-stage Peltier cooler was used to maintain a substrate temperature of -15°C.

5 Deposition conditions are shown in Table 3.

Table 3. Sputtering conditions for Pd metallization on mercuric iodide.

Ar Pressure	Power	Deposition Rate	Substrate Temperature	Target-Substrate Spacing	Deposition Time	Base Pressure
5.0 mTorr	100 W D.C.	700 Å/min	- 15 °C	8.0 cm	2.0 min	< 1 x 10 ⁻⁶ Torr

Lithography for Pd patterning on HgI₂

Shadow masks are typically used to pattern Pd (or other materials such as gold and platinum) on HgI₂ because typical lithography processes present two major problems. First, acetone and/or other solvents are used to remove the photoresist after patterning. Many solvents, including acetone, however, attack HgI₂ rapidly. Second, HgI₂ cannot withstand the high temperatures typically required for photoresist baking. Shadow masks, however, are quite limiting in terms of minimum pattern size and freedom of design. Fine line widths of guard electrodes are required for electron-only operation of HgI₂ spectrometers, and shadow masks are often not suitable for generating them.

As a result, a new lithography process was developed which is compatible with HgI₂. A low-temperature photoresist (requiring a moisture-removal bake of only 50°C) was employed in the new process. Second, the required use of standard solvents for dissolving the photoresist after metal etching was circumvented. This was achieved through two steps. First, a high-temperature bake, which typically polymerizes a photoresist and makes it impervious to a developer solution, was not used. The palladium etchant was chosen such that a soft-bake was sufficient to make the photoresist a suitable mask for the Pd etch step. Without this hardbake step, the photoresist removal can subsequently be

promoted by a flood exposure step after the Pd etching. The flood exposure makes the photoresist soluble in the standard developer solution of 2:1 TMAH:H₂O, which does not attack mercuric iodide. The processing steps and materials used are shown in Table 4.

5 **Table 4.** Lithography steps for Pd metallization patterning

Step	Parameters	Chemical
Spin adhesion promoter on HgI ₂	3000 rpm, 20 sec	HMDS
Spin photoresist on HgI ₂	3000 rpm, 20 sec, 1.3 μm thick	OCG 825
Soft-bake	90°C, 2min, (50°C max temp reached)	
Mask alignment, exposure	200 W Hg lamp, 1.8 min exposure	
Develop photoresist	50 sec, H ₂ O rinse, N ₂ dry	2:1 TMAH: H ₂ O
Soft-bake	90°C, 2min	
Pd etch	1 min	2:10:30 Br:HNO ₃ :HCl
Flood exposure	200 W Hg lamp, 2.5 min exposure	
Photoresist removal	2 min, H ₂ O rinse, N ₂ dry	2:1 TMAH: H ₂ O

Sublimation

Trenches were etched by a sublimation process after the first Pd metallization and patterning steps. The vapor pressure of HgI₂ below its structural transformation temperature of 129°C is less than 11 mTorr. Therefore, sublimation should be governed by temperature only and not by gas flow as long as the pressure is kept somewhat below 11 mTorr. As a result, a single zone quartz furnace with liquid nitrogen vapor trap that was turbo-pumped to a pressure of 3 mTorr was sufficient for trench etching. The temperature outside of the furnace tube (but inside the heater housing) was monitored and the average

10

15

temperature in the furnace over a 20 minute interval was estimated to be approximately 95-120°C.

In addition to being temperature dependent, the sublimation rate was also found to be dependent on the particular etchant used for the Pd patterning, with an increase in Br content generating higher etch rates and a redder surface. (One possible explanation is that Br in the Pd etchant solution prevents the mercuric iodide from losing excessive iodine and thus making the surface more red. Because iodine has a higher vapor pressure than mercury under these conditions, an increased sublimation rate is obtained.) Without Br, the surface was dark (typical of Hg-rich surfaces) and it sublimed at a slower rate.

After the sublimation step was completed, the entire surface was metallized with Pd again and then patterned with collection and grid electrodes having the dimensions shown in Table 5. Photographs of a fabricated device are shown at various stages of the fabrication process in Figures 1A-1E. FIGURE 1A shows lithography on Pd to define a grid etch pit. FIGURE 1B shows etching of Pd. FIGURE 1C shows HgI₂ sublimation etch (300 µm). FIGURE 1D shows lithography for grid and collecting electrodes. FIGURE 1E shows mercuric iodide after Pd etching.

Figure 2 shows the basic structure of a coplanar grid gamma-ray detector 10 of the present invention. HgI₂ crystal 12 has been fabricated with trenches 14. Electrode material 16 (e.g., palladium, gold or platinum) has been deposited into the trenches as well as onto the adjacent outer wall 18 and the opposite wall 20. The detector is electrically biased such that the electrodes on the outer wall 18 are at the most positive potential, and the electrodes within the trenches 14 are at a less positive potential, while the electrode on the wall 20 is at a potential that is negative with respect to the potential at wall 18 and within trenches 14. A capacitor 22 and resistor 24 in series may be used to isolate the electrodes on wall 18 from the electrodes within trenches 14. In operation, radiation (such as a gamma photon) creates electron-hole pairs. The holes drift in the direction of the negative potential at wall 20 and the electrons drift to the positive potentials at

the trenches and the more positive potential at the outer wall **18**. The detector signal is read only from electrons passing between the electrodes in trenches **14** and the electrodes on the outer wall **18** within the measurement region **26**. The wires may be attached to each set of said metal layers using material carbon

5 Aquadag or another colloidal suspension of graphite particles. Aquadag is a trade name for a colloidal suspension of graphite in water, generally used as a lubricant and a conductive coating.

Table 5. Dimensions used for fabrication of HgI₂ electron-only spectrometer.

Collection width	Grid width	Collection length	Pit width	Pit depth
600 μm	400 μm	6.0 mm	1000 μm	300 μm

10 The present invention is technically different in both the equipment required and the potential achievable performance. Standard lithography equipment is required, because it provides the capability to achieve smaller electrode feature sizes and electrodes that are more closely spaced. Etched trench devices are not currently realized in commercial HgI₂ spectrometers and

15 thus the sublimation process provides a new and different capability to fabricate such devices.

The lithography is an improvement because (as an optical method) much finer feature size can be obtained than with commonly used shadow mask techniques, allowing for enhanced ability to localize gamma-ray interactions, engineer the internal electric field in the devices, and generate higher quality

20 images of shaped radiation sources.

The sublimation adds additional design freedom and consequently performance. Currently there are no commercial methods for generating fine-line cross-strip imaging arrays, fine-line coplanar grids, small-pixels having a small

25 pitch, or etched trench detectors of HgI₂. Implementation of the new processing steps leads to x-ray and gamma-ray spectrometers with both improved spatial imaging capability and energy resolution.

Industrial suppliers will use this new process for fabricating HgI₂ x-ray and gamma-ray and charged-particle spectrometers for medical imaging, environmental monitoring, baggage handling, and materials sorting applications. The government will use HgI₂ detectors and instruments utilizing these sensors to detect nuclear smuggling, safeguard dismantled nuclear materials, search for nuclear weapons, verify treaty compliance, locate unexploded ordnance, detect contraband drugs, perform forensic analyses, conduct x-ray astronomy, cleanup waste sites, and act as personal dosimeters.

The primary cost of HgI₂ radiation detectors is in the crystal growth and secondarily fabrication. Standard lithography equipment, evacuated furnaces, and sputtering equipment are required for the current process. It is estimated that these additional fabrication systems would represent less than 20% of total production cost.

The foregoing description of the invention has been presented for purposes of illustration and description and is not intended to be exhaustive or to limit the invention to the precise form disclosed. Many modifications and variations are possible in light of the above teaching. The embodiments were chosen and described to best explain the principles of the invention and its practical application to thereby enable others skilled in the art to best use the invention in various embodiments and with various modifications suited to the particular use contemplated. The scope of the invention is to be defined by the following claims.

CLAIMS

What is claimed is:

1. A method of lithographically patterning a metal deposited onto a HgI_2 crystal,
5 comprising the steps of:

depositing a metal layer onto a HgI_2 crystal surface;

spinning adhesion promoter onto said metal layer;

spinning photoresist onto said adhesion promoter;

soft-baking said photoresist to remove moisture;

10 aligning a mask to said photoresist;

exposing a portion of said photoresist that is not concealed by said mask
to produce exposed photoresist and unexposed photoresist;

removing said mask from said photoresist;

developing and removing said exposed photoresist;

15 soft-baking said photoresist a second time;

etching said metal layer;

flood exposing said unexposed photoresist to produce a second exposed
photoresist; and

developing and removing said second exposed photoresist.

20

2. The method of claim 1, wherein said metal layer comprises metal selected
from the group consisting of palladium, gold, carbon, indium-tin-oxide, chromium,
tantalum and platinum.

25 3. The method of claim 1, wherein the step of spinning adhesion promoter is
carried out with HMDS at about 3000 rpm for about 20 seconds.

4. The method of claim 1, wherein the step of spinning photoresist is carried out
with OCG 825 at about 3000 rpm for about 20 seconds or until said photoresist is
30 about 1.3 μm thick.

5. The method of claim 1, wherein the step of soft-baking said photoresist to remove moisture is carried out at about 90°C ambient temperature for about 2 minutes, wherein said photoresist temperature does not exceed 50°C.

5 6. The method of claim 1, wherein the step of exposing said photoresist is carried out at about 200 W with a Hg lamp at an exposure time of about 1.8 minutes.

10 7. The method of claim 1, wherein the step of developing said photoresist is carried out with a mixture of about 2:1 TMAH: H₂O for about 50 seconds followed by rinsing with H₂O and drying with N₂.

15 8. The method of claim 1, wherein said metal layer comprises palladium, wherein the step of etching said metal layer is carried out with a mixture of about 2:10:30 of Br:HNO₃:HCl for about 1 minute.

20 9. The method of claim 1, wherein the step of flood exposing said unexposed photoresist is carried out with about 200 W from a Hg lamp at an exposure time of about 2.5 minutes.

10. The method of claim 1, wherein the step of developing and removing said second exposed photoresist is carried out with a mixture of about 2:1 of TMAH: H₂O for about 2 minutes followed by rinsing with H₂O and drying with N₂.

25 11. The method of claim 1, wherein the step of developing and removing said second exposed photoresist will produce uncovered HgI₂, the method further comprising subliming said uncovered HgI₂ to produce trenches.

12. The method of claim 11, wherein said HgI_2 crystal is held at a vacuum pressure that is less than 11 mTorr, wherein sublimation is governed by temperature of the HgI_2 surface.

- 5 13. The method of claim 12, wherein said vacuum pressure is about 3 mTorr and said temperature is within a range of about 95-120°C.

14. The method of claim 11, further comprising depositing metal into said trenches.

10

15. The method of claim 1, further comprising smoothing said HgI_2 crystal surface prior to the step of depositing a metal layer.

15

16. The method of claim 15, wherein the step of smoothing said HgI_2 crystal surface further comprises:

placing said HgI_2 crystal into a solution of $\text{KI}:\text{H}_2\text{O}$; and
agitating said solution.

20

17. The method of claim 16, wherein said $\text{KI}:\text{H}_2\text{O}$ is at a mixture of about 1:10.

18. The method of claim 16, where the KI in said solution is about 5% to 15% of said solution.

25

19. The method of claim 16, further comprising agitating said solution at an agitation rate within a range from about 100 rpm to about 200 rpm.

20. The method of claim 16, wherein the step of agitating said solution comprises (i) agitating said solution in a mixture of about 1:10 of KI:H₂O at about 200 rpm with a magnetic stir bar for about 30 seconds at each orientation of 0°, 180°, 270° and 90°, (ii) rinsing said crystal with H₂O and (iii) drying said crystal with N₂.

21. The method of claim 1, wherein the step of depositing a metal layer comprises sputtering metal onto said HgI₂ crystal surface.

22. The method of claim 1, wherein the step of depositing metal is carried out with a metal deposition technique selected from the group consisting of metal sputtering and metal evaporation.

23. The method of claim 22, wherein said HgI₂ crystal is placed in a vacuum prior to metal deposition, wherein said vacuum comprises a low base pressure.

24. The method of claim 22, wherein said HgI₂ crystal is cooled to promote metal adhesion.

25. The method of claim 24, wherein said HgI₂ crystal is cooled to a temperature of about -15°C.

26. The method of claim 25, wherein said HgI₂ crystal is cooled with a two-stage Peltier cooler.

27. The method of claim 23, wherein said low base pressure is less than about 1×10^{-6} T.

28. The method of claim 21, wherein the step of sputtering metal onto said Hgl₂ surface further comprises:

placing said Hgl₂ crystal into a vacuum chamber having a base pressure of less than about 1×10^{-6} T;

- 5 cooling said Hgl₂ crystal to about -15 °C;
 filling said vacuum chamber with a working gas for sputtering; and
 depositing metal onto said Hgl₂ crystal surface.

29. The method of claim 28, wherein said working gas for sputtering comprises
10 Ar at about 5 mTorr.

30. The method of claim 28, wherein said metal is deposited at a rate of about 700 Å/minute for about 2 minutes.

15 31. The method of claim 28, wherein the step of depositing metal comprises depositing palladium.

32. The method of claim 1, further comprising attaching wires to said metal layer, wherein said wires are for applying electrical voltages and providing electrical
20 connection to signal conditioning electronics.

33. The method of claim 32, wherein said wires are attached to each set of said metal layers using material selected from the group consisting of carbon aquadag and colloidal suspension of graphite particles.

25

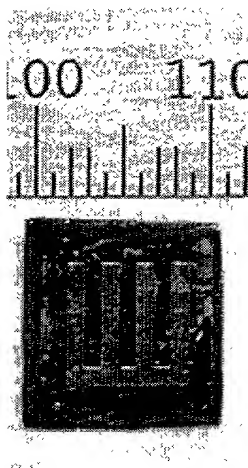
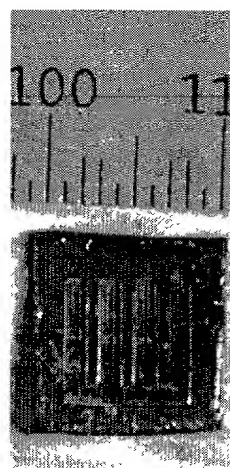
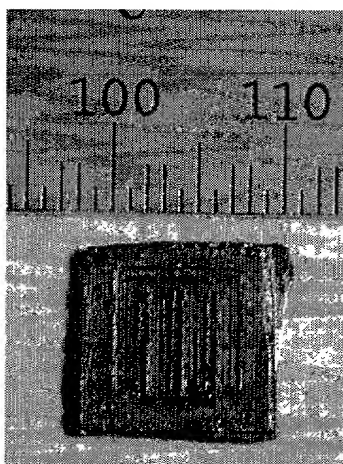
34. The method of claim 1, further comprising depositing insulating material onto said Hgl₂ crystal surface and said metal layer.

35. The method of claim 34, wherein said insulating material is parylene C.

30

ABSTRACT OF THE DISCLOSURE

- A photolithographic process forms patterns on HgI_2 surfaces and defines metal sublimation masks and electrodes to substantially improve device performance by increasing the realizable design space. Techniques for smoothing HgI_2 surfaces and for producing trenches in HgI_2 are provided. A sublimation process is described which produces etched-trench devices with enhanced electron-transport-only behavior
- 5

FIG. 1A**FIG. 1B****FIG. 1C****FIG. 1D****FIG. 1E****FIG. 1A-1E**

Mercuric iodide after various processing steps: (1A) lithography on Pd to define grid etch pit. (1B) etching of Pd. (1C) Hgl_2 sublimation etch ($300\ \mu\text{m}$). (1D) lithography for grid and collecting electrodes. (1E) after Pd etching.

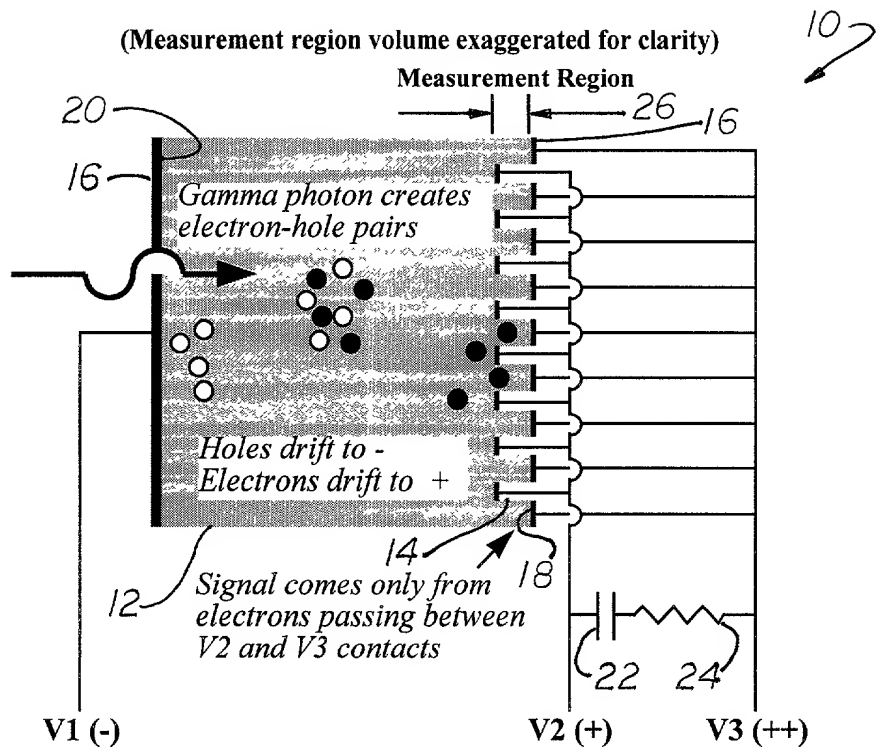


FIG. 2

DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name.

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled:

LITHOGRAPHY PROCESS FOR PATTERNING HgI₂ PHOTONIC DEVICES

the specification of which (check one) X is attached hereto, or was filed on , as application Serial No. and was amended on (if applicable).

I hereby state that I have reviewed and understand the contents of the identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, §1.56.

I hereby claim the benefit under Title 35, United States Code, §119(e) of any United States provisional application(s) listed below.

Application Number	Filing Date	Status
--------------------	-------------	--------

Application Number	Filing Date	Status
--------------------	-------------	--------

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, §1.56 (a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

Application Number	Filing Date	Status
--------------------	-------------	--------

POWER OF ATTORNEY I hereby appoint the following attorneys and practitioners to prosecute this application and to transact all business in the United States Patent Office connected therewith:

George H. Libman, Registration No. 27,984; Gregory A. Cone, Registration No. 28,835; Kurt C. Olsen, Registration No. 29,961; Timothy D. Stanley, Registration No. 30,428; and Timothy P. Evans, Registration No. 41,013.

CUSTOMER BAR CODE

Send correspondence to: Timothy Evans,
Sandia National Laboratories
Post Office Box 969, Mail Stop 9031
Livermore, CA 94551-0969

Direct telephone calls to: Timothy Evans at (925) 294-3690

10/31/00 TUE 07:56 FAX 617 258 4238

DRAPER LAB E20

002

Attorney Docket SD-8310

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under §1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Full name of first inventor: Mark J. MESCHER
Inventor's signature: *Mark J. Mescher* Date: Oct. 29, 2000
Residence: 2053 Commonwealth Avenue, Auburndale, MA 02466
Citizenship: USA
Post Office Address: 2053 Commonwealth Avenue, Auburndale, MA 02466

Full name of first inventor: Ralph B. JAMES
Inventor's signature: _____ Date: _____
Residence: 5420 Lenore Avenue, Livermore, CA 94550
Citizenship: USA
Post Office Address: 5420 Lenore Avenue, Livermore, CA 94550

Full name of first inventor: Haim HERMON
Inventor's signature: _____ Date: _____
Residence: 2 Bar Kochva Street, Jerusalem, State of Israel
Citizenship: USA
Post Office Address: 2 Bar Kochva Street, Jerusalem, State of Israel

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under §1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Full name of first inventor: Mark J. MESCHER
 Inventor's signature: _____ Date: _____
 Residence: 2053 Commonwealth Avenue, Auburndale, MA 02466
 Citizenship: USA
 Post Office Address: 2053 Commonwealth Avenue, Auburndale, MA 02466

Full name of first inventor: Ralph B. JAMES
 Inventor's signature: *Ralph B. James* Date: Nov. 9, 2000
 Residence: 5420 Lenore Avenue, Livermore, CA 94550
 Citizenship: USA
 Post Office Address: 5420 Lenore Avenue, Livermore, CA 94550

Full name of first inventor: Haim HERMON
 Inventor's signature: _____ Date: _____
 Residence: 2 Bar Kochva Street, Jerusalem, State of Israel
 Citizenship: USA
 Post Office Address: 2 Bar Kochva Street, Jerusalem, State of Israel

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under §1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

Full name of first inventor: Mark J. MESCHER
 Inventor's signature: _____ Date: _____
 Residence: 2053 Commonwealth Avenue, Auburndale, MA 02466
 Citizenship: USA
 Post Office Address: 2053 Commonwealth Avenue, Auburndale, MA 02466

Full name of first inventor: Ralph B. JAMES
 Inventor's signature: _____ Date: _____
 Residence: 5420 Lenore Avenue, Livermore, CA 94550
 Citizenship: USA
 Post Office Address: 5420 Lenore Avenue, Livermore, CA 94550

Full name of first inventor: Haim HERMON
 Inventor's signature: Haim Hermon Date: Nov. 1, 00
 Residence: 2 Bar Kochva Street, Jerusalem, State of Israel
 Citizenship: USA
 Post Office Address: 2 Bar Kochva Street, Jerusalem, State of Israel